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                 Web Page URLs for STN Seminar Schedule - N. America
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NEWS
                  "Ask CAS" for self-help around the clock
NEWS
         OCT 23
                 The Derwent World Patents Index suite of databases on STN
                 has been enhanced and reloaded
                 CHEMLIST enhanced with new search and display field
NEWS
         OCT 30
NEWS
         NOV 03
                 JAPIO enhanced with IPC 8 features and functionality
NEWS
         NOV 10
                 CA/CAplus F-Term thesaurus enhanced
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         NOV 10
                 STN Express with Discover! free maintenance release Version
                  8.01c now available
NEWS
      8
         NOV 20
                 CA/CAplus to MARPAT accession number crossover limit increased
                 to 50,000
NEWS 9
         DEC 01
                 CAS REGISTRY updated with new ambiguity codes
NEWS 10
         DEC 11
                 CAS REGISTRY chemical nomenclature enhanced
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                 WPIDS/WPINDEX/WPIX manual codes updated
NEWS 12
         DEC 14
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NEWS 13
         DEC 18
                 CA/CAplus pre-1967 chemical substance index entries enhanced
                  with preparation role
NEWS 14
         DEC 18
                 CA/CAplus patent kind codes updated
NEWS 15
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                 MARPAT to CA/CAplus accession number crossover limit increased
                  to 50,000
NEWS 16
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                 MEDLINE updated in preparation for 2007 reload
NEWS 17
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NEWS 18
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NEWS 19
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NEWS 20
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                 WPIDS/WPINDEX/WPIX enhanced with IPC 8 reclassification data
NEWS 21
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         JAN 22
NEWS 24
         JAN 29
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NEWS 25
         JAN 29
                 CAS Registry Number crossover limit increased to 300,000 in
                 multiple databases
NEWS 26
         FEB 13
                 CASREACT coverage to be extended
                 PATDPASPC enhanced with Drug Approval numbers
NEWS 27
         Feb 15
NEWS 28
         Feb 15 RUSSIAPAT enhanced with pre-1994 records
NEWS 29
         Feb 23
                 KOREAPAT enhanced with IPC 8 features and functionality
NEWS 30
         Feb 26 MEDLINE reloaded with enhancements
NEWS 31
         Feb 26 EMBASE enhanced with Clinical Trial Number field
NEWS 32
         Feb 26
                 TOXCENTER enhanced with reloaded MEDLINE
NEWS 33
         Feb 26
                 IFICDB/IFIPAT/IFIUDB reloaded with enhancements
NEWS 34
         Feb 26
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                 to 300,000 in multiple databases
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NEWS EXPRESS NOVEMBER 10 CURRENT WINDOWS VERSION IS V8.01c, CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP), AND CURRENT DISCOVER FILE IS DATED 25 SEPTEMBER 2006.

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FULL ESTIMATED COST

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http://www.cas.org/infopolicy.html

=> s formoterol

944 FORMOTEROL

1 FORMOTEROLS

L1944 FORMOTEROL

(FORMOTEROL OR FORMOTEROLS)

=> s 73573-87-2

REG1stRY INITIATED

Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

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=> s steroid
       112480 STEROID
       113670 STEROIDS
       171188 STEROID
                (STEROID OR STEROIDS)
=> s L4 and L5
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=> dup rem L6
PROCESSING COMPLETED FOR L6
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L8
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     1529 FLUTICASONE
L10
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L11
=> d 1-38 ibib abs L9
    ANSWER 1 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
                      2006:666025 CAPLUS
DOCUMENT NUMBER:
                       145:152690
TITLE:
                       Method for inducing crystalline state transition in
                       pharmaceuticals
INVENTOR(S):
                       Nakamichi, Kouichi; Izumi, Shougo; Oka, Masaaki
PATENT ASSIGNEE(S):
                       Nippon Shinyaju Company, Ltd., Japan
                       U.S., 18 pp., Cont.-in-part of U.S. 5,456,923.
SOURCE:
                       CODEN: USXXAM
DOCUMENT TYPE:
                       Patent
LANGUAGE:
                       English
FAMILY ACC. NUM. COUNT: 2
PATENT INFORMATION:
    PATENT NO.
                       KIND DATE
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                                                               DATE
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    US 5811547
                                         US 1995-416815
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                              19980922
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    CA 2147279
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    AU 9351607
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                              19950802
                                                                19931013 <--
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                        В1
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                                        AT 1993-922625
                                                                19931013 <--
    ES 2145063
                                          ES 1993-922625
                        Т3
                              20000701
                                                                19931013 <--
    US 5456923
                                         US 1993-129133
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Α

PRIORITY APPLN. INFO.:

19951010

JP 1992-303085

WO 1993-JP1469 US 1993-129133 19931115 <--

A 19921014 <--W 19931013 <--

A2 19931115 <--

JP 1991-112554 A 19910416 <--WO 1992-JP470 W 19920414 <--

AB This invention has for its object to provide a method of inducing a transition in crystalline state of a crystallizable pharmaceutical with great ease and improved efficiency and uniformity on a high production scale. An extruder is used for inducing a transition from one crystalline state ( $\Delta$ ) to another crystalline state in a crystallizable pharmaceutical. An extruded indomethacin (form  $\alpha$ ) was converted to an amorphous form.

REFERENCE COUNT: 10 THERE ARE 10 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 2 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2003:355834 CAPLUS

DOCUMENT NUMBER: 138:362665

TITLE: Immunostimulatory nucleic acids for the treatment of

asthma and allergy

INVENTOR(S): Bratzler, Robert L.; Petersen, Deanna M.; Fouron, Yves

PATENT ASSIGNEE(S): USA

SOURCE: U.S. Pat. Appl. Publ., 221 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
US 2003087848	A1	20030508	US 2001-776479	20010202 <
US 2004067902	A9	20040408		
US 2004235774	A1	20041125	US 2004-831778	20040423 <
US 2006154890	A1	20060713	US 2005-301360	20051209 <
US 2007037767	A1	20070215	US 2006-526896	20060922 <
PRIORITY APPLN. INFO.:			US 2000-179991P	P 20000203 <
			US 2001-7,76479	A1 20010202 <
			US 2004-831778	A1 20040423
			US 2005-301360	A1 20051209

OTHER SOURCE(S): MARPAT 138:362665

AB The invention involves administration of an immunostimulatory nucleic acid alone or in combination with an asthma/allergy medicament for the treatment or prevention of asthma and allergy in subjects. The combination of drugs are administered in synergistic amts. or in various dosages or at various time schedules. The invention also relates to kits and compns. concerning the combination of drugs.

L9 ANSWER 3 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2003:44146 CAPLUS

DOCUMENT NUMBER:

138:73178

TITLE:

Preparation and pharmaceutical combinations of [(hetero)arylalkyl]piperidinyl amine, amide, or carbamate CCR3 antagonists for treatment of asthma,

allergic disease, or inflammation

INVENTOR(S):

Bahl, Ash; Perry, Matthew; Springthorpe, Brian

PATENT ASSIGNEE(S): Astrazeneca AB, Swed.

SOURCE:

Brit. UK Pat. Appl., 91 pp.

CODEN: BAXXDU

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
GB 2373186 PRIORITY APPLN. INFO.:	A	20020918	GB 2001-4534 GB 2001-4534	20010223 < 20010223 <

OTHER SOURCE(S): MARPAT 138:73178 GΙ

$$R^{1} - (Q)_{m} - (CR^{2}R^{3})_{n} - T - \begin{pmatrix} X^{2} - X^{1} \\ N - Z - R^{6} \\ X^{3} - X^{4} \end{pmatrix}$$

$$\begin{array}{c|c} F & & \\ \hline \\ N & \\ H & \\ \end{array}$$

AB Title compds. I [wherein Z = CR4R5, CO, or CR4R5Z1; Z1 = alkylene, alkenylene, or CONH; R1 = (un)substituted alkyl, alkenyl, (hetero)cycloalkyl, or (hetero)aryl; Q = O, S, NR9, CO, CONR9, NR9CO, or CH=CH; m = 0-1; n = 0-6 with the proviso that when n = 0; then m = 0; R2 and R3 = independently H or alkyl; or CR2R3 = (alkyl)cycloalkyl; T = NR10, CONR10, NR11CONR10, or CONR10R11; X1-X4 = independently CH2CHR12 or CO; R4 and R5 = independently H or alkyl; R6 = (un) substituted (hetero) aryl; R9-R11 = independently H, alkyl, haloalkyl, hydroxyalkyl, cycloalkyl(alkyl), or phenylalkyl; R12 = independently (cyclo)alkyl or CO; or R12 groups of X1 and X3 or X4, or X2 and X3 or X4 join to form CH2CH2, CH2CH2CH2, CH2OCH2, or CH2SCH2; or pharmaceutically acceptable salts or solvates thereof] were prepared as cysteine-cysteine chemokine receptor 3 (CCR3) antagonists for use in pharmaceutical combinations with a histamine antagonist, steroid, leukotriene modulator, human cytokine,  $\beta$ -agonist, phosphodiesterase inhibitor, or antibody (no data). example, 1-(3,4-dichlorobenzyl)-4-piperidinamine • 2CF3CO2H was condensed with 2-(4-fluorophenyl) acetic acid to give N-[1-(3,4dichlorobenzyl)-4-piperidinyl]-2-(4-fluorophenyl)acetamide (II). useful in combination therapy for the treatment of asthma, rhinitis, and other allergic or inflammatory conditions (no data).

ANSWER 4 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2002:832576 CAPLUS

DOCUMENT NUMBER: 137:346197

TITLE: Treatment of respiratory and lung diseases with

antisense oligonucleotides and a bronchodilating agent

INVENTOR(S): Nyce, Jonathan W.; Li, Yukui; Sandrasagra, Anthony;

Katz, Evan; Pabalan, Jonathan; Aguilar, Douglas;

Miller, Shoreh; Tang, Lei; Shahabuddin, Syed

PATENT ASSIGNEE(S): Epigenesis Pharmaceuticals, Inc., USA

SOURCE: PCT Int. Appl., 764 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT NO	•		KIN	D :	DATE		i	APPL	ICAT:	ION I	.00		D	ATE	
				-											
WO 200208	5309		A2		2002.	T03T	1	WO 21	002-	JSI3.	143		21	J0204	423 <
W: A	E, AG,	AL,	AM,	AT,	ΑU,	AZ,	BA,	BB,	BG,	BR,	BY,	ΒZ,	CA,	CH,	CN,
C	O, CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	ES,	FI,	GB,	GD,	GE,	GH,
G	M, HR,	ΗU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	ΚP,	KR,	ΚZ,	LC,	LK,	LR,
	S, LT,														
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UA, UG, US, UZ, VN, YU, ZA, ZM, ZW RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG US 2004049022 US 2003-627930 A1 20040311 20030725 PRIORITY APPLN. INFO .: US 2001-286036P P 20010424 <--WO 2002-US13135 A2 20020423 WO 2002-US13143 A2 20020423

MARPAT 137:346197 OTHER SOURCE(S):

This patent relates to a composition comprising a carrier, oligonucleotides (oligos) that are antisense to adenosine receptors, and contain low amts. of or no adenosine (A), plus bronchodilating agents. All antisense oligonucleotides designed in accordance with the invention were highly effective at countering or reducing effects mediated by the receptors to which they are targeted. Two antisense phosphorothioated oligos targeting human adenosine A1 receptor mRNA, one targeting adenosine A2b receptor, and two targeting an A3 receptor are capable of countering the effect of exogenously administered adenosine which is mediated by the specific receptor they are targeted to. The activity of the antisense oligos are specific to the target and substitutively fail to inhibit another target. An oligonucleotide wherein the phosphodiester bonds are substituted with phosphorothioate bonds evidenced an unexpected superiority over the phosphodiester antisense oligo. In addition, they result in extremely low or non-existent deleterious side effects or toxicity. This represents 100% success in providing agents that are highly effective and specific in the treatment of bronchoconstriction and/or inflammation. These agents and the composition and formulations provided are suitable for the treatment of respiratory tract, pulmonary and malignant diseases associated with bronchoconstriction, respiratory tract inflammation and allergies, impaired airways, including lung disease and diseases whose secondary effects afflict the lungs of a subject, such as allergies, asthma, impeded respiration, allergic rhinitis, pain, cystic fibrosis, pulmonary fibrosis, RDA, COPD, and cancers, among others. The present agents and composition may be administered preventatively, prophylactically or therapeutically in conjunction with other therapies, or may be utilized as a substitute for therapies that have significant, neg. side effects. The method of the present invention is also practiced with antisense oligonucleotides targeted to many genes, mRNAs and their corresponding proteins in essential the same manner.

ANSWER 5 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2002:832564 CAPLUS

DOCUMENT NUMBER: 137:329451

TITLE: Pharmaceutical formulations and kit for treatment of

respiratory and lung disease with non-glucocorticoid

steroids and/or ubiquinone and a

bronchodilating agent

INVENTOR(S):

Nyce, Jonathan W.

PATENT ASSIGNEE(S):

Epigenesis Pharmaceuticals, Inc., USA

PCT Int. Appl., 49 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

SOURCE:

English

FAMILY ACC. NUM. COUNT:

PATENT	NO.			KIN	) 1	DATE			APPL	ICAT:	ION I	NO.		Di	ATE	
	D 2002085296 A2 20021031 D 2002085296 A3 20030403 W: AE, AG, AL, AM, AT, AU, AZ,						Ţ	WO 2	002-	US12	552		2	0020	122 <	
W:	AE, CO, GM, LS,	CR, HR,	AL, CU, HU,	AM, CZ, ID,	AT, DE, IL,	AU, DK, IN,	AZ, DM, IS,	DZ, JP,	EC, KE,	EE, KG,	ES, KP,	FI, KR,	GB, KZ,	GD, LC,	GE, LK,	GH, LR,

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PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
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     US 2003216329
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                                20031120
                                           US 2003-461563
     US 2005070487
                         Α1
                                20050331
                                           US 2004-475689
                                                                  20040812 <--
PRIORITY APPLN. INFO .:
                                           US 2001-286139P
                                                               P 20010424 <--
                                           WO 2002-US12552
                                                               A1 20020422
                                           US 2002-388170P
                                                               P 20020612
OTHER SOURCE(S):
                        MARPAT 137:329451
     A pharmaceutical or veterinary composition, comprises a first active agent
     selected from a non-glucocorticoid steroid or analogs, a
     ubiquinone, or salts thereof, and a second active agent comprising a
     bronchodilator. The composition is provided in various formulations and in the
     form of a kit. The products of this patent are applied to the prophylaxis
     and treatment of respiratory, lung and malignant diseases.
    ANSWER 6 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
                         2002:813911 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         137:316082
TITLE:
                         Formoterol/steroid bronchodilating
                         compositions and methods of use thereof
INVENTOR(S):
                         Banerjee, Partha S.; Chaudry, Imitiaz A.
PATENT ASSIGNEE(S):
                         Dey LP, USA
                         PCT Int. Appl., 52 pp.
SOURCE:
                         CODEN: PIXXD2
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
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                               DATE
                                                                  DATE
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                         A2
                               20021024
                                          WO 2002-US6252
                                                                  20020301 <--
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                         A3
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            GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
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                                                                 20010622 <--
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     EP 1385494
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                                          EP 2002-719098
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            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
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AB Bronchodilating compns. intended for administration as a nebulized aerosol are provided. In certain embodiments, the compns. contain formoterol, or a derivative thereof, and a steroidal anti-inflammatory agent. Methods for treatment, prevention, or amelioration of one or more symptoms of bronchoconstrictive disorders using the compns. provided herein are also provided. For example, a solution was prepared containing Formoterol fumarate dihydrate 85  $\mu \rm g/mL$ , budesonide 125

JP 2002-580917

US 2002-145978

US 2001-284607P

US 2001-887496

WO 2002-US6252

20020301 <--

20020513 <--

P 20010417 <--

A1 20010622 <--

W 20020301

IE, SI, LT, LV, FI, RO, MK, CY, AL, TR

Т

A1

20050512

20021205

JP 2005512944

US 2002183293

PRIORITY APPLN. INFO.:

μg/mL, vitamin E TPGS 10 μg/mL, Polyethylene glycol 10 μg/mL, citrate buffer 50mM, sodium chloride 7.5 mg/mL, and water as needed.

ANSWER 7 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2002:730354 CAPLUS

DOCUMENT NUMBER: 137:252996

Ipratropium formulation for pulmonary inhalation TITLE: INVENTOR(S):

Wulffhart, Harold; Ayoub, Khaldoun; Logiudice,

Rosemary; Piskorz, Hanna

Pharmascience, Can. PATENT ASSIGNEE(S):

U.S., 9 pp. SOURCE: CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATĖ	APPLICATION NO.	DATE
US 6455028	B1	20020924	US 2001-841181	20010423 <
CA 2441549	A1	20021031	CA 2002-2441549	20020403 <
WO 2002085338	A2	20021031	WO 2002-CA450	20020403 <
WO 2002085338	A3	20030403		
WO 2002085338	B1	20030703		
W: AU, CA,	JP			
RW: AT, BE,	CH, CY, D	E, DK, ES,	FI, FR, GB, GR, IE,	IT, LU, MC, NL,
PT, SE,			•	
EP 1381353	A2	20040121	EP 2002-713970	20020403 <
R: AT, BE,	CH, DE, DI	K, ES, FR,	GB, GR, IT, LI, LU,	NL, SE, MC, PT,
IE, FI,				, ,
PRIORITY APPLN. INFO	. :		US 2001-841181	A 20010423 <

WO 2002-CA450 W 20020403 AB Pharmaceutical aerosol formulations are provided comprising substantially nonacicular particles of a bronchodilator selected from the group consisting of ipratropium and pharmacol. acceptable salts, solvates, hydrates, esters and isomers thereof. The described formulations include a propellant selected from the group consisting of a fluorocarbon propellant, a hydrogen-containing fluorocarbon propellant, and mixts. thereof. The formulations are substantially free of both surfactant and solvent. Methods of use and drug delivery devices are also provided. For example, an ipratropium bromide inhaler was prepared using approx. 5.5 mg of ipratropium bromide nonacicular particles and 8.3 g of 1,1,1,2-tetrafluoroethane (HFC-134a). Upon repeated actuation, the

inhaler delivered about 20 µg of the active agent per dose without clogging. REFERENCE COUNT: 23 THERE ARE 23 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 8 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2002:555336 CAPLUS

DOCUMENT NUMBER: 137:114526

TITLE: A method for the preparation of nanoparticles INVENTOR(S): Watanabe, Wiwik; Kauppinen, Esko; Ahonen, Petri;

Brown, David; Muttonen, Esa

PATENT ASSIGNEE(S): Orion Corporation, Finland SOURCE: PCT Int. Appl., 22 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE --------------

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WO 2002056866
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             GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
             LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
             PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
             UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU,
             TJ, TM
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             CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
             BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
    EP 1351666
                         A1
                               20031015
                                         EP 2002-710900
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            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
                                          JP 2002-557374
     JP 2004520157
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                               20040708
                                                                  20020118 <--
     US 2004091542
                         Α1
                               20040513
                                           US 2003-466365
                                                                  20031211 <--
PRIORITY APPLN. INFO.:
                                           FI 2001-115
                                                               A 20010118 <--
                                           WO 2002-FI42
                                                               W 20020118
    The invention relates to free nano-sized particles of active agents e.g.
     therapeutic, cosmetic or diagnostic agents, and to a method for the preparation
    of such particles. The method comprises providing a liquid feed stock
     comprising an active agent or combination of two or more active agents,
     atomizing the liquid feed stock, suspending the droplets in a carrier gas,
    and passing the carrier gas and droplets through a heated tube flow
     reactor under predetd. residence time and temperature history, and collecting
    the particles produced. Nano-sized crystalline spherical uncharged particles
    with narrow aerodynamic particle size distribution and rough surfaces, are
     obtained. The particles show improved dissoln. rate in-vitro and
    bioavailability in-vivo, dispersibility and stability. Nanosized
    beclomethasone dipropionate particles were prepared
REFERENCE COUNT:
                        4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS
                              RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 9 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
                        2002:449474 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                        137:11011
TITLE:
                        Particulate inhalation carriers
                        Buckton, Graham; Al-Hadithi, Dima; Brocchini, Stephen
INVENTOR(S):
PATENT ASSIGNEE(S):
                        School of Pharmacy, University of London, UK
SOURCE:
                        PCT Int. Appl., 21 pp.
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
LANGUAGE:
                        English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                         APPLICATION NO.
    PATENT NO.
                        KIND
                               DATE
                                                                 DATE
                                          -----
                               20020613 WO 2001-GB5436
    WO 2002045682
                        A1
                                                                 20011210 <--
        W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
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            GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
            LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH,
            PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TN, TR, TT, TZ,
            UA, UG, US, UZ, VN, YU, ZA, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU,
            TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH,
            CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
            BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
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20020618

IE, SI, LT, LV, FI, RO, MK, CY, AL, TR

20030903

Α5

Α1

AU 2002022145

EP 1339388

JP 2004517834

AU 2002-22145

AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,

20040617 JP 2002-547468

EP 2001-999355

20011210 <--

20011210 <--

US 2003-433435 20031020 <--GB 2000-30074 A 20001208 <--WO 2001-GB5436 W 20011210 <--US 2004062719 A1 20040401 PRIORITY APPLN. INFO.: The present invention provides a particulate substrate suitable for carrying a drug for delivery, comprising a substantially crystalline core and a surface coating, wherein the particulate substrate has a proportion of amorphous character of 2% or greater by weight of particulate substrate, and a process for the production of carrier particles comprising the steps of: (a) mixing crystalline particles having an average diameter greater than 10 µm with at least partially amorphous particles having average diams. less than 10 µm; (b) exposing the mixture to conditions capable of inducing crystallization of the amorphous particles for a predetd. period in order that partial crystallization takes place. The core material is selected from saccharides, most preferably lactose and the surface of the substrate is formed from the same material as the core. The drug is selected from steroids, hormones, therapeutic proteins and peptides,  $\beta$ -2 agonists, bronchodilators, corticosteroids and antihistamines. REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 10 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2002:51237 CAPLUS DOCUMENT NUMBER: 136:123631 TITLE: Aerosol formulation containing a polar fluorinated compound INVENTOR(S): Rogueda, Philippe Astrazeneca AB, Swed. PATENT ASSIGNEE(S): PCT Int. Appl., 61 pp. SOURCE: CODEN: PIXXD2 DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE
WO 2002003958 A1 20020117 WO 2001-SE1606 20010710 <--W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NÒ, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG 20020117 CA 2001-2415092 20010710 <-- 20030423 EP 2001-952071 20010710 <--CA 2415092 Α1 EP 1303258 Α1 EP 1303258 20061011 В1 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR

US 2003-332568 20030109 <-NO 2003-133 20030110 <-GB 2000-16876 A 20000711 <-WO 2001-SE1606 W 20010710 <--The present invention relates to a stable pharmaceutical aerosol AB formulation intended for inhalation. The formulation contains an active

BR 2001-12322

JP 2002-508413

NZ 2001-523379

AT 2001-952071

ZA 2003-75

20010710 <--

20010710 <--

20010710 <--

20010710 <--

20030103 <--

20030708

BR 2001012322 A JP 2004502719 T

PRIORITY APPLN. INFO.:

JP 2004502719 T 20040129
NZ 523379 A 20040625
AT 342048 T 20061115
ZA 2003000075 A 20040405
US 2003194378 A1 20031016
NO 2003000133 A 20030224
RITY APPLN. INFO.:

substance, an aerosol propellant, a polar fluorinated mol. and an excipient. The preferred propellant is HFA 134a or HFA 227 or a mixture Thus, an aerosol formulation contained budesonide 0.125, methoxy-PEG-DSPE 0.320, 1H,1H,2H,2H-perfluorooctan-1-ol 31.7 and HFA-227 to 100%. REFERENCE COUNT: THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 11 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:867964 CAPLUS

DOCUMENT NUMBER: 135:376803

TITLE: Stable pharmaceutical solution formulations for

pressurized metered dose inhalers

INVENTOR(S): Lewis, David; Ganderton, David; Meakin, Brian;

Brambilla, Gaetano; Ferraris, Alessandra

Chiesi Farmaceutici S.P.A., Italy PATENT ASSIGNEE(S):

SOURCE:

Eur. Pat. Appl., 18 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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PATENT NO.
                      KIND DATE APPLICATION NO.
                                                              DATE
                      A1 20011128 EP 2001-112230 20010518 <--
    EP 1157689
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                                         CA 2000-2411047
    CA 2411047
                              20011129
                       A1
                                                               20000522 <--
    WO 2001089480
                              20011129
                                       WO 2000-EP4635
                       Α1
                                                               20000522 <--
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            ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU,
            LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE,
            SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW
        RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
            DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
            CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                    A
    BR 2000015884
                            20030708 BR 2000-15884
                                                               20000522 <--
                                        HU 2003-2007
    HU 200302007
                       A2
                              20030929
                                                               20000522 <--
                      T
    JP 2003534266
                              20031118
                                        JP 2001-585725
                                                              20000522 <--
    EE 200200649
                      Α
                              20040615
                                        EE 2002-649
                                                               20000522 <--
                       A2
                                        EP 2004-11423
    EP 1466594
                              20041013
                                                               20010518 <--
    EP 1466594
                             20041201
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    BG 107256
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                                                               20021108 <--
    NO .2002005568
                                         NO 2002-5568
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    HK 1058900
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                        Α1
                              20060127
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PRIORITY APPLN. INFO.:
                                                           A 20000522 <--
                                         WO 2000-EP4635
                                         EP 2001-112230
                                                           A3 20010518 <--
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An aerosol solution composition for use in an aerosol inhaler comprises an active

material, a propellant containing a hydrofluoroalkane, a cosolvent and optionally a low volatility component to increase the mass median aerodynamic diameter (MMAD) of the aerosol particles on actuation of the inhaler. The active ingredient is a  $\beta 2$  agonist selected from salbutamol, formoterol, salmeterol, and TA-2005, salts thereof or their combination with steroid such as beclomethasone dipropionate, fluticasone propionate, budesonide, and its 22R-epimer or an anticholinergic atropine-like derivative such as ipratropium bromide, oxitropium bromide, and tiotropium bromide. The composition is stabilized by using a small amount of mineral acid and a suitable can having part or all of its internal metallic surfaces made of stainless steel, anodized aluminum or lined with an inert organic coating.

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 12 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:833060 CAPLUS

DOCUMENT NUMBER: 135:376741

TITLE: Stable metal ion-lipid powdered pharmaceutical

compositions

INVENTOR(S):
Dellamary, Luis A.; Riess, Jean; Schutt, Ernest G.;

Weers, Jeffry G.; Tarara, Thomas E. Alliance Pharmaceutical Corp., USA

SOURCE: PCT Int. Appl., 42 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 14

PATENT INFORMATION:

PATENT ASSIGNEE(S):

PATENT NO. KIND DATE								1	APPL	ICAT:	I NOI	. OV		D.	ATE			
WO.	2001	0851	37		7.2	-	2001	1115	•		001-1				- 2	0010	500	/
	2001						2001			NO 2	001-	JS140	324		۷	0010.	300	
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BY,	BZ,	CA,	CH,	CN,	
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		HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KΡ,	KR,	ΚZ,	LC,	LK,	LR,	LS,	LT,	
				-				MN,										
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		YŲ,	ZA,	ZW														
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	2408							1115										
EP	1282	405			A2		2003	0212		EP 2	001-	9331	94		2	0010	508	<
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	2003							1111										
	2006							0316								0060		
	2006				A1		2006	1207								0061		
PRIORIT	Y APP	LN.	INFO	. :							000-					0000		
											999-1					9990		
											001-					0010		
									1	WO 2	001-	JS148	324		W 2	0010	508	<

AB Microparticle compns. comprising metal ion-lipid complexes for drug delivery are described including methods of making the microparticle compns. and methods of treating certain conditions and disease states by administering the microparticle compns. The metal ion-lipid complexes can be combined with various drugs or active agents for therapeutic administration. The microparticle compns. of the present invention have superior stability to other microparticle compns. resulting in a microparticle composition with longer shelf life and improved dispersibility. The microparticle compns. of the present invention have a transition temperature

(Tm) of at least 20° above the recommended storage temperature (Tst) for drug delivery. An aqueous preparation was prepared by mixing two prepns., A and B,

immediately prior to spray drying. The preparation A was comprised of a fluorocarbon-in-water emulsion in which 26 g perfluorocctyl bromide was dispersed in 33 g water with the aid of 1.30 g of SPC-3 emulsifier (hydrogenated soy phosphatidylcholine). The preparation B contained 0.162 g CaCl2.2H20 and 0.162 g budesonide dissolved/suspended in 4 g water. The resulting microparticle of the sample had a PL-budesonide-CaCl2.2H20 weight ratio of about 80:10:10. The mean volume aerodynamic particle size of the dry powder was approx. 4.1  $\mu m$ .

L9 ANSWER 13 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:659249 CAPLUS

DOCUMENT NUMBER: 135:366235

TITLE: Is there a need for another inhalative  $\beta$ 2-agonist

besides formoterol in patients with asthma?

AUTHOR(S): Matthys, Heinrich

CORPORATE SOURCE: Medizinische Klinik, Abteilung Pneumologie

Universitatsklinik Freiburg, Freiburg, Germany

SOURCE: Respiration (2001), 68(4), 432-437

CODEN: RESPBD; ISSN: 0025-7931

PUBLISHER: S. Karger AG

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

AB A review with refs. Formoterol can substitute the rapid- and short-acting  $\beta 2$ -agonists as well as the slow- and long-acting salmeterol. Therefore formoterol in a fixed combination with an inhalant steroid reduces the aerosol devices necessary for asthma control to only one, to be used for regular "controller" and, as needed, "rescue therapy". The side effect profile of formoterol is comparable to the short-acting  $\beta 2$ -agonists which makes the combination with a topically active glucocorticoid applicable in patients of any asthma severity as long as they are able to perform an inspiratory

vital capacity maneuver.

REFERENCE COUNT:

63

THERE ARE 63 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 14 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:548882 CAPLUS

DOCUMENT NUMBER:

136:63413

TITLE:

Long-acting  $\beta$ -agonists and steroids -

trial experience

AUTHOR(S):

Lofdahl, C.-G.

CORPORATE SOURCE:

Department of Respiratory Medicine, Lund University

Hospital, Lund, SE-22185, Swed.

SOURCE:

Clinical & Experimental Allergy Reviews (2001

), 1(1), 18-22

CODEN: CEARC3; ISSN: 1472-9725

PUBLISHER: Blackwell Science Ltd.
DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

AB A review on the effect of a combination of a low-dose inhaled steroid and  $\beta$ 2-agonists, salmeterol or formoterol,

on clin. outcomes in asthma. This combined treatment system improves the lung function symptom control and the quality of life in patients with persistent asthma to a greater extent than increasing the dose of the inhaled steroid. However, increasing the maintenance dose of the inhaled steroid might be more appropriate and effective

the inhaled steroid might be more appropriate and effective treatment strategy in preventing repeated severe exacerbations in asthma patients.

REFERENCE COUNT:

THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 15 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:475821 CAPLUS

DOCUMENT NUMBER:

CORPORATE SOURCE:

135:267426

TITLE:

Effect of oral prednisolone on the bronchoprotective

effect of formoterol in patients with

persistent asthma

AUTHOR(S):

Grootendorst, D. C.; Sterk, P. J.; Heijerman, H. G. M. Dept of Pulmonology, Leijenburg Hospital, The Hague,

NL-2504 LN, Neth.

SOURCE:

European Respiratory Journal (2001), 17(3),

374-379

CODEN: ERJOEI; ISSN: 0903-1936 European Respiratory Society

PUBLISHER: DOCUMENT TYPE: Journal LANGUAGE: English

Tolerance to the bronchoprotective effects by long-acting  $\beta$ 2-agonists (LAB) in patients with asthma is not prevented by inhaled corticosteroids (ICS). This study examined whether oral prednisolone can restore the bronchoprotective effects of formoterol in 24 patients with persistent asthma already treated with ICS (at least 800 μg budesonide day-1 or equivalent) and LAB, using a parallel-group design. During a 2-wk run-in period and during the study, patients used formoterol 12 µg twice daily by Turbuhaler, instead of their own LAB. At baseline and at the end of 7-days treatment with oral placebo or prednisolone (30 mg·day-1), provocative concentration of histamine causing a 20% fall in forced expiratory volume in one second (PC20 histamine) was measured on two sep. days after randomized singledose inhalation of placebo (postP) or formoterol (postF). In addition, PC20postF was measured 24 h after starting oral treatment. The protective effect by formoterol at baseline and during treatment was calculated as the difference between the logs of PC20postP and PC20postF. The mean±SEM in doubling dose (DD) bronchoprotective effect at baseline was 0.8±0.4 DD in the placebo group and  $1.0\pm0.4$  DD in the prednisolone group. At the end of the treatment period, the protective effect changed to 1.0±0.5 DD and 0.8±0.6 DD in the placebo and prednisolone treated groups, resp. This change was not different between the groups (p>0.4). In conclusion, the bronchoprotective effect by formoterol is not influenced by 1 wk prednisolone treatment in patients with asthma who are using regular inhaled corticosteroids and long-acting  $\beta$ 2-agonists. These findings indicate that tolerance to long-acting \$2-agonists cannot be restored by oral steroid therapy.

REFERENCE COUNT: THERE ARE 25 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 16 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2001:475820 CAPLUS

DOCUMENT NUMBER:

135:266981

TITLE:

Reversing acute bronchoconstriction in asthma: The effect of bronchodilator tolerance after treatment

with formoterol

AUTHOR(S):

Jones, S. L.; Cowan, J. O.; Flannery, E. M.; Hancox,

R. J.; Herbison, G. P.; Taylor, D. R.

CORPORATE SOURCE:

Depts of Medical and Surgical Sciences, Dunedin School

of Medicine, University of Otago, Dunedin, N. Z.

SOURCE:

European Respiratory Journal (2001), 17(3),

368-373

CODEN: ERJOEI; ISSN: 0903-1936 European Respiratory Society

DOCUMENT TYPE:

Journal English

LANGUAGE:

PUBLISHER:

Continuous treatment with a short-acting  $\beta$ 2-agonist can lead to

reduced bronchodilator responsiveness during acute bronchoconstriction. This study evaluated bronchodilator tolerance to salbutamol following regular treatment with a long-acting  $\beta2$ -agonist, formoterol The modifying effect of i.v. corticosteroid was also studied. asthmatic subjects (using inhaled steroids) participated in a randomized, double-blind, placebo-controlled, cross-over study. Formoterol 12  $\mu$ g b.i.d. or matching placebo was given for 10-14 days with > 2 wk washout. Following each treatment, patients underwent a methacholine challenge to induce a fall in forced expired volume in one second (FEV1) of at least 20%, then salbutamol 100  $\mu g$ , 100  $\mu g$ , and 200  $\mu g$  was inhaled via a spacer at 5 min intervals, with a further 400 μg at 45 min. Following each treatment, patients underwent a methacholine challenge to induce a fall in forced expired volume in one second (FEV1) of at least 20%, then salbutamol 100  $\mu g$ , 100  $\mu g$ , and

200  $\mu$ g was inhaled via a spacer at 5 min intervals, with a further 400  $\mu$ g at 45 min. After a third single-blind formoterol treatment period, hydrocortisone 200 mg was given i.v. prior to salbutamol. Dose-response curves for change in FEV1 with salbutamol were compared using anal. of covariance to take account of methacholine-induced changes in spirometry. Regular formoterol resulted in a significantly lower FEV1 after salbutamol at each time point compared to placebo (p<0.01). The area under the curves (AUCs) for 15 (AUC0-15) and 45 (AUC0-45) min were 28.8% and 29.5% lower following formoterol treatment (p<0.001). Pretreatment with hydrocortisone had no significant modifying effect within 2 h of administration. It is concluded that significant tolerance to the bronchodilator effects of inhaled salbutamol occurs 36 h after stopping the regular administration of formoterol. This bronchodilator tolerance is evident in circumstances of acute bronchoconstriction.

REFERENCE COUNT: 33 THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 17 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001:360974 CAPLUS

DOCUMENT NUMBER: 135:235689

Long-acting β2-agonists

AUTHOR(S):

Johnson, Malcolm; Hagan, Gerry W. E.

CORPORATE SOURCE:

GlaxoWellcome, Uxbridge, UK

SOURCE:

TITLE:

Progress in Respiratory Research (2001),

31 (New Drugs for Asthma, Allergy and COPD), 60-63

CODEN: PRRRAE; ISSN: 1422-2140

PUBLISHER:

S. Karger AG

DOCUMENT TYPE:

Journal; General Review

LANGUAGE: English

AB A review with 36 refs. The LABAs salmeterol and formoterol have both prolonged airway smooth muscle effects and non-bronchodilator activity. They have a complementary mode of action to the topical anti-inflammatory effects of corticosteroids, and inhibit mucosal edema, increase mucociliary transport and reduce respiratory tract infection. In asthma, LABAs are currently positioned as "add-on" therapy, where combination with inhaled steroids results in better lung function and symptom control, decreased rescue medication and fewer exacerbations. In COPD patients, LABAs such as salmeterol reduce breathlessness, decrease exacerbations and improve health-related quality of life.

REFERENCE COUNT: 36 THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 18 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2001

2001:247172 CAPLUS

DOCUMENT NUMBER:

134:256899

TITLE:

Combination of loteprednol and  $\beta 2\text{-adrenoceptor}$ 

agonists for the treatment of allergies and

respiratory tract diseases

INVENTOR(S):

Szelenyi, Istvan; Poppe, Hildegard; Heer, Sabine;

Engel, Juergen

PATENT ASSIGNEE(S):

Asta Medica Ag, Germany PCT Int. Appl., 16 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent German

LANGUAGE:

SOURCE:

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FAMILY ACC. NUM. COUNT:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2001022956	A2	20010405	WO 2000-EP9392	20000926 <
WO 2001022956	A3	20011011		

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             TR, UA, US, UZ, YU, ZA, AM, AZ, MD, TJ, TM
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     BR 2000014374
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                                 20020625
                                             BR 2000-14374
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     EP 1216047
                          A2
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PRIORITY APPLN. INFO.:
                                             DE 1999-19947235
                                                                  A 19990930 <--
                                             WO 2000-EP9392
                                                                  W 20000926 <--
     The invention relates to a novel combination of a soft steroid,
     especially loteprednol, and at least one \beta2-adrenoceptor agonist for
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treating allergies and/or respiratory tract diseases simultaneously, sequentially or sep.; to drugs containing said combination, to methods for producing such drugs and to the use of the novel combination for producing drugs for the simultaneous, sequential or sep. treatment of allergies and/or respiratory tract diseases. Thus and aerosol was prepared that contained 6  $\mu$ g formoterol fumarate dihydrate and 200  $\mu$ g loteprednol per stroke. 2H-heptafluoropropane (1.000 g) propellant was cooled to  $-55\,^{\circ}\text{C}$  and 11.7 g Tagat TO in 11.7 g ethanol was added under stirring, followed by the addition of 3.34 g micronized loteprednol etabonate and 0.1 g formoterol fumarate dihydrate. The suspension was diluted with 1,170.0 g 2H-heptafluoropropane, filled in metal containers with valves for dosing 50  $\mu$ L suspension per stroke.

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ANSWER 19 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
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ACCESSION NUMBER: 2001:152458 CAPLUS

DOCUMENT NUMBER: 134:183526

TITLE: Method to produce powders for pulmonary or nasal

administration

INVENTOR(S): Woolfe, Austen John; Zeng, Xian Ming; Langford, Alan

PATENT ASSIGNEE(S): Norton Healthcare Ltd., UK SOURCE: PCT Int. Appl., 18 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATEN	T NO.			KIN	D	DATE			APPL:	ICAT	ION :	NO.		Di	ATE	
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CA 23	82216			A1		2001	0301		CA 2	-000	2382	216		2	0000	821 <

JP 2003526629 Т 20030909 JP 2001-518024 20000821 <--US 1999-150095P P 19990820 <--WO 2000-GB3230 W 20000821 <--PRIORITY APPLN. INFO.: A pharmaceutical formulation comprises a mixture of two or more drugs optionally together with one or more excipients, the mixture being formed by the steps of: co-crystallization or co-precipitation of the drugs followed by micronization or milling to produce a uniform powder having a particle size and other properties suitable for formulation for pulmonary or nasal administration. An aqueous solution of 5% salbutamol sulfate:ipratropium bromide (10:1) mixture was prepared and was spray dried. The diameter of particles was less than 3  $\mu$ m. REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 20 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN 2001:63851 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 134:120962 Powders consisting of particles with a perfectly TITLE: smooth surface, for use as carriers for the preparation of inhalation mixtures with micronized drugs and method for their preparation Caponetti, Giovanni; Catellani, Pier Luigi; Bettini, INVENTOR(S): Ruggero; Colombo, Paolo; Ventura, Paolo Chiesi Farmaceutici S.p.A., Italy PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 39 pp. CODEN: PIXXD2 DOCUMENT TYPE: Patent English LANGUAGE: FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION: KIND DATE APPLICATION NO. PATENT NO. DATE WO 2001005429 A2 20010125 WO 2000-EP6690 20000713 <--WO 2001005429 A3 20011004 W: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG IT 99MI1582 20010116 IT 1999-MI1582 19990716 <--Α1 EP 1196146 20020417 EP 2000-956180 20000713 <--Α2 EP 1196146 В1 20060913 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, CY Α BR 2000012351 20020611 BR 2000-12351 20000713 <--AT 339191 T 20061015 AT 2000-956180 20000713 <--B1 A1 US 6780508 20040824 US 2002-30686 20020416 <--US 2005118113 20050602 US 2004-806240 20040323 <--

AB Carriers for use in the preparation of mixts. for inhalation powders intended for pulmonary administration of micronized drugs by means of a dry powder inhaler and the method for their preparation are described. An inhalation powder of beclometasone dipropionate mixed with smoother  $\alpha$ -lactose monohydrate carrier was prepared

IT 1999-MI1582

WO 2000-EP6690 US 2002-30686 A 19990716 <--W 20000713 <--

A1 20020416

L9 ANSWER 21 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2000:645829 CAPLUS

PRIORITY APPLN. INFO.:

DOCUMENT NUMBER: 133:227824

TITLE: Modified carrier particles for use in dry powder

inhalers

INVENTOR(S):
Musa, Rossella; Bilzi, Roberto; Ventura, Paolo;

Chiesi, Paolo

PATENT ASSIGNEE(S): Chiesi Farmaceutici S.P.A, Italy

SOURCE: PCT Int. Appl., 25 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

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AB The invention relates to carrier particles for use in pharmaceutical compns. for the pulmonary administration of medicaments by means of dry powder inhalers. In particular, the invention relates to a novel technol. process for obtaining a carrier modified so as to improve the efficiency of redispersion of active particles and hence increase the respirable fraction. After the treatment of the invention, the surface of said modified carrier particles can also be coated with a suitable additive so as to further improve the respirable fraction.  $\alpha$ -Lactose monohydrate 99.75 % was mixed with 0.25% magnesium stearate and 200  $\mu$ g/dose beclomethasone-17,21-dipropionate. The flowability properties of the carrier did not change significantly even in the presence of ternary mixture and a significant increase of the fine particle fraction was observed with the carrier.

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 22 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2000:645828 CAPLUS

DOCUMENT NUMBER: 133:227823

TITLE: Improved powdery pharmaceutical compositions for

inhalation comprising low percentage lubricant Musa, Rossella; Ventura, Paolo; Chiesi, Paolo

Chiesi Farmaceutici S.P.A., Italy

PATENT ASSIGNEE(S):

SOURCE:

PCT Int. Appl., 36 pp. CODEN: PIXXD2

DOCUMENT TYPE:

INVENTOR(S):

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE										ION :			Di	ATE			
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EP	1158	958			A1		2001	1205		EP 1	999-	9155	47		1	9990	305 <
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US	6528	096			В1		2003	0304		US 2	000-	6036	20		20	0000	626 <
US	2003	13388	30		A1		2003	0717		US 2	003-	3581	74		20	0030	205 <
US	2006	2573	30		A1		2006	1116		US 2	006-	4921	05		20	0060	725 <
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							:			US 2	000-	6036	20		A3 2	0000	626 <- <b>-</b>
										US 2	003-	3581	74	7	A1 2	0030	205

AΒ The invention describes the use of a little percentage of lubricant (0.05-0.5%) by weight) in powdery pharmaceutical compns. for use in dry powder inhalers in order to increase the fine particle dose. A process for coating the surface of the carrier particles with such little amount of lubricant is also claimed. The use of limited amount of the lubricant is safe and allows to prepare ordered stable mixts. without segregation of the active particles during handling and before use.  $\alpha ext{-Lactose}$ monohydrate was mixed with 0.1%, 0.25%, or 0.5% magnesium stearate and 100, 200, and 400 µg/dose beclomethasone-17,21-dipropionate. Multidose devices filled with the mixture were then tested. No significant increase in fine particle dose was obtained from the concentration of magnesium stearate above 0.25%.

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 23 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2000:277843 CAPLUS

DOCUMENT NUMBER:

132:313698

TITLE:

Storable active substance concentrate with

formoterol

INVENTOR(S):

Hochrainer, Dieter; Zierenberg, Bernd

PATENT ASSIGNEE(S):

Boehringer Ingelheim Pharma K.-G., Germany

SOURCE: PCT Int. Appl., 17 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE -----

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PRIORITY APPLN. INFO.:
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                                                                W 19991009 <--
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AB A storage-stable concentrate of the antiasthmatic, formoterol, in the form of a solution or suspension for use in inhalers contains formoterol base or a salt or addition product thereof at a concentration of 75-500 mg formoterol/mL in a polar (preferably protic) liquid, e.g. aqueous NaCl solution, EtOH, or a mixture thereof. The formulation may addnl.

contain an inorg. or organic acid to adjust the pH to 2.0-7.0, preservatives, antioxidants, complexing agents, and addnl. active substances such as  $\beta\text{-mimetics}$ , cholinergic antagonists, antiallergic agents, leukotriene antagonists, and/or steroids. Thus, a concentrate comprised 5 mg formoterol (particle size 5  $\mu\text{m}$ ) in 0.015 mL 20 weight% aqueous NaCl solution adjusted to pH 5.0 with fumaric acid. An inhalant was prepared by mixing this suspension with 4.5 mL H2O/EtOH (1:1) containing benzalkonium chloride 0.45 and Na EDTA 2.25 mg and adjusting the pH to 5.0 with HCl.

L9 ANSWER 24 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2000:254113 CAPLUS

DOCUMENT NUMBER: 132:284231

TITLE: Storable formulation of active substance INVENTOR(S): Hochrainer, Dieter; Zierenberg, Bernd

PATENT ASSIGNEE(S): Boehringer Ingelheim Pharma K.-G., Germany

SOURCE: Ger. Offen., 8 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 4

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                                                                  A3 19991009 <--
                                             WO 1999-EP7589
                                                                    19991009 <--
                                                                  W
                                             US 1999-416476
                                                                  A1 19991012 <--
                                             US 2001-871500
                                                                  A1 20010531 <--
                                             US 2002-256781
                                                                  A1 20020927
AB
     A storage-stable formulation of an active substance in the form of a
```

AB A storage-stable formulation of an active substance in the form of a concentrated solution or suspension in an atomizer or cartridge is provided for use

in inhalers. The concentrate is diluted with H2O or solvent immediately before the 1st use of the composition. Stability of suspended particles of the active substance in the formulation is enhanced by addition of an alkali metal or ammonium chloride or salt of an organic acid. The active substance may be a  $\beta$ -mimetic, anticholinergic, or antiallergic drug, platelet-activating factor antagonist, leukotriene antagonist, and/or steroid. Thus, a suspension of 5 mg formoterol (particle size 5  $\mu m$ ) in 0.015 mL water was adjusted to pH 5.0 with fumaric acid for storage. This suspension was diluted with 4.5 mL H2O/EtOH (1:1) containing benzalkonium chloride 0.45 and Na EDTA 2.25 mg, adjusted to pH 5.0 with HCl, for inhalation.

ANSWER 25 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN 2000:254112 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 132:284230 TITLE: Storable liquid formoterol formulation INVENTOR(S): Hochrainer, Dieter; Zierenberg, Bernd PATENT ASSIGNEE(S): Boehringer Ingelheim Pharma K.-G., Germany Ger. Offen., 8 pp. CODEN: GWXXBX DOCUMENT TYPE: Patent LANGUAGE: German FAMILY ACC. NUM. COUNT: 4 PATENT INFORMATION: APPLICATION NO. PATENT NO. KIND DATE DATE ----\_\_\_\_\_ A1 20000420 DE 1998-19847969 19981017 <--A1 20000427 CA 1999-2343123 19991009 <--DE 19847969 20000427 CA 1999-2343123 CA 2343123 19991009 <--A2 20000427 WO 1999-EP7581 A3 20000803 19991009 <--WO 2000023065 WO 2000023065 W: AU, BG, BR, CA, CN, CZ, EE, HR, HU, ID, IL, IN, JP, KR, LT, LV, MX, NO, NZ, PL, RO, SG, SI, SK, TR, UA, US, UZ, VN, YU, ZA, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE AU 1999-62019 AU 9962019 Α 20000508 19991009 <--AU 764126 В2 20030814 BR 9914507 Α 20010626 BR 1999-14507 19991009 <--20010808 EP 1999-948972 EP 1121112 A2 19991009 <--EP 1121112 B1 20020605 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO TR 200101096 T2 20011221 TR 2001-200101096 19991009 <--20020529 HU 2001-3925 HU 200103925 A2 19991009 <--20020615 AT 1999-948972 20020617 EE 2001-224 AT 218331 T 19991009 <--EE 200100224 A 20020617 EE 2001-224

EE 4219 B1 20040216

JP 2002527473 T 20020827 JP 2000-576840

JP 3636430 B2 20050406

PT 1121112 T 20021129 PT 1999-948972

ES 2178479 T3 20021216 ES 1999-948972

NZ 511225 A 20030829 NZ 1999-511225

SK 285382 B6 20061207 SK 2001-494

US 6150418 A 20001121 US 1999-416474

TW 562676 B 20031121 TW 1999-88117879

BG 105391 A 20011130 BG 2001-105391

NO 2001001663 A 20010403 NO 2001-1663

HR 2001000255 A1 20020430 HR 2001-255

ZA 2001003056 A 20020123 ZA 2001-3056

HK 1041448 A1 20050513 HK 2002-103088

JP 2005047933 A 20050930 IN 2005-MN468

RITY APPLN. INFO.: DE 1998-19847969 EE 200100224 A 19991009 <--19991009 <--19991009 <--19991009 <--19991009 <--19991009 <--19991012 <--19991015 <---20010329 <--20010403 <--20010406 <--

W 19991009 <--A3 20010322 <--IN 2001-MN321 A storage-stable formulation of the  $\beta$ 2-adrenergic agonist, formoterol, in the form of a concentrated solution or suspension is provided for use in inhalers for inhalational or nasal therapy of asthma. The concentrate is diluted with H2O or solvent immediately before the 1st use of

PRIORITY APPLN. INFO.:

JP 2004-323973

US 1998-112380 US 1998-112380P

JP 2000-576840

WO 1999-EP7581

DE 1998-19847969 A 19981017 <--

20010412 <--20020424 <--

20041108 <--20050520 <--

P 19981214 <--P 19981214 <--

A3 19991009 <--

the composition Stability of the formulation is enhanced by addition of an organic

or inorg. acid, preferably in combination with a complexing agent, especially when the solvent contains EtOH. The formulation may also contain addnl.  $\beta\text{-mimetics}$ , anticholinergics, antiallergic drugs, platelet-activating factor antagonists, leukotriene antagonists, and/or steroids. Thus, a suspension of 5 mg formoterol (particle size 5  $\mu\text{m}$ ) in 0.015 mL 20 weight% NaCl was adjusted to pH 5.0 with fumaric acid for storage. This suspension was diluted with 4.5 mL H2O/EtOH (1:1) containing benzalkonium chloride 0.45 and Na EDTA 2.25 mg, adjusted to pH 5.0 with HCl, for inhalation.

REFERENCE COUNT:

THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 26 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2000:92868 CAPLUS

DOCUMENT NUMBER: 132:117313

TITLE: Bronchodilator response to albuterol after regular

formoterol and effects of acute corticosteroid

administration

AUTHOR(S): Lipworth, Brian J.; Aziz, Imran

CORPORATE SOURCE: Department of Clinical Pharmacology and Therapeutics

and Respiratory Medicine Ninewells Hospital and

Medical School, University of Dundee, Dundee, DD1 9SY,

UK

22

SOURCE: Chest (2000), 117(1), 156-162

CODEN: CHETBF; ISSN: 0012-3692

PUBLISHER: American College of Chest Physicians

DOCUMENT TYPE: Journal LANGUAGE: English

There is controversy about the development of bronchodilator subsensitivity after regular administration of long-acting  $\beta$ 2-agonists. The purpose of the study was to evaluate whether regular treatment with formoterol affects the bronchodilator response to repeated puffs of albuterol, and also to assess the effects of acute administration of a bolus dose of IV or inhaled corticosteroid. Twelve patients (mean [SD] age, 43 [15] years; FEV1, 57 [17] % predicted) with stable, moderate to severe persistent asthma who were all taking inhaled corticosteroids were evaluated in a randomized, placebo-controlled, double-blind, double-dummy, crossover study. Patients received treatments each for 2 wk followed by a bolus (IV/inhaled) of corticosteroid or placebo: (1) placebo inhaler bid + bolus placebo; (2) formoterol Turbuhaler 24  $\mu g$  metered dosage bid (delivered dosage 18  $\mu$ g bid) + placebo; (3) formoterol 24  $\mu$ g bid + bolus IV hydrocortisone, 200 mg; or (4) formoterol 24  $\mu$ g bid + bolus inhaled budesonide,  $1,600~\mu g$ . Bronchodilator response to repeated puffs of albuterol (200 to 1,600  $\mu$ g) for > 80 min was measured at 2 h after bolus administration of placebo or corticosteroid. The study was powered at the 80% level to detect a 20% difference in area under curve between 20 and 80 min (AUC) for FEV1 response to albuterol as change from baseline (primary end point). There was significant subsensitivity (p = 0.01) of the mean albuterol FEV1 response (as AUC, L + s) after formoterol alone (737) as compared to placebo (1,453) along with partial reversal by steroid administration: formoterol + hydrocortisone (1,050), and formoterol + budesonide (942). There was a similar pattern of subsensitivity (p = 0.03) for the mean albuterol forced expiratory flow between 25% and 75% of vital capacity response (as AUC, L): placebo (2,149), formoterol alone (1,002), formoterol + hydrocortisone (1,402), and formoterol + budesonide (1,271). Regular treatment with formoterol produced significant bronchodilator subsensitivity to repeated puffs of albuterol, which was partially reversed by a bolus dose of systemic or inhaled corticosteroid.

REFERENCE COUNT:

THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 27 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2000:15933 CAPLUS

DOCUMENT NUMBER: 132:45179

TITLE: Asthma quality of life during 1 year of treatment with

budesonide with or without formoterol

AUTHOR(S): Juniper, E. F.; Svensson, K.; O'Byrne, P. M.; Barnes, P. J.; Bauer, C-A; Lofdahl, C-G, A; Postma, D, S;

P. J.; Bauer, C-A.; Lofdahl, C-G. A.; Postma, D. S.;

Pauwels, R. A.; Tattersfield, A. E.; Ullman, A. CORPORATE SOURCE: Dept of Clinical Epidemiology & Biostatistics,

McMaster University, Hamilton, ON, Can. European Respiratory Journal (1999), 14(5),

1038-1043

CODEN: ERJOEI; ISSN: 0903-1936

PUBLISHER: Munksgaard International Publishers Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

SOURCE:

AB The Formoterol and Corticosteroids Establishing Therapy (FACET)

study has provided the first opportunity to examine the long-term effects of inhaled steroids and long-acting  $\beta 2$ -agonists on

asthma-specific quality of life. The objectives of the present study were

to: evaluate the effects of long-term (1 yr) formoterol and increasing doses of budesonide on asthma quality of life; 2) to determine whether initial improvements in quality of life are sustained when

improvements in clin. indexes persist; and 3) to evaluate the long-term relationship between changes in clin. indexes and changes in quality of life. Of the 852 asthmatic adults enrolled, 470 from five countries participated in this quality of life evaluation. After a 4-wk run-in on

1,600  $\mu$ g budesonide, patients were randomized to either 200  $\mu$ g (Bud200) or 800  $\mu$ g budesonide (Bud800) in combination with either 24

µg formoterol (F) or placebo daily for 1 yr. The Asthma Quality of Life Questionnaire (AQLQ) was completed and conventional clin. indexes measured at enrollment and randomization and on seven occasions during the following 12 mo. During the run-in, there was an improvement

in AQLQ score (changes (∆) in overall score≈0.50; p<0.0001).

After randomization, there was a further improvement in the Budge

After randomization, there was a further improvement in the Bud800+F group ( $\Delta$ =0.21; p=0.028). One month post-randomization, improvements in all groups stabilized and were sustained throughout the 12 mo in a pattern very similar to that observed for the conventional clin. indexes. The correlation of individual patient changes in clin. indexes and changes in AQLQ score during the 12-mo randomized period were weak to moderate (maximum r=0.51). Improvements in quality of life, which were greatest in the 800

 $\mu g$  budesonide plus 24  $\mu g$   $\,$  formoterol group, were sustained throughout the 12 mo in a similar manner to the clin. indexes. Long-term changes in conventional clin. indexes cannot be used to predict the effect

of treatment on individual patient experience.

REFERENCE COUNT: 15 THERE ARE 15 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 28 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1999:133202 CAPLUS

DOCUMENT NUMBER: 130:200925

TITLE: Finely divided pharmaceutical particles for inhalation

INVENTOR(S): Briggner, Lars-Erik; Bystrom, Katarina; Jakupovic,

Edib; Trofast, Eva; Trofast, Jan

PATENT ASSIGNEE(S): Astra Aktiebolag, Swed.

SOURCE: U.S., 7 pp., Cont.-in-part of U.S. Ser. No. 459,660.

CODEN: USXXAM

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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A 19990223 US 1996-606655
      US 5874063
                                                                                     19960226 <--
                            A 19921117
B2 19950907
      AU 9215347
                                        19921117 AU 1992-15347
                                                                                    19920324 <--
      AU 662519
      EP 580648
                                                      EP 1992-907877
                                A1
                                        19940202
                                                                                     19920324 <--
      EP 580648
                                В1
                                       19960508
           R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, MC, NL, SE
      JP 06506454 T
                                     19940721
                                                      JP 1992-507195
                                                                                    19920324 <--
      JP 3400999
                                B2
                                         20030428
      EP 680752
                               A2
                                        19951108
                                                      EP 1995-111178
                                                                                    19920324 <--
      EP 680752
                               А3
                                        19951122
      EP 680752
                               B1
                                        20011114
    PL 168232
RU 2112507
SK 280310
B6 19991108
CZ 286936
B6 20000816
CZ 1993-2116
JP 2003155228
A 20030527
JP 2002-347368
NO 9303575
A 19931006
NO 1993-3575
NO 311867
B1 20020211
FI 105388
B1 20000815
FI 1993-4429
US 5709884
A 19980120
US 1995-379471
US 5637620
A 19970610
US 1995-459660
A 19961008
US 1995-479494
SE 1991-1090
SE 1993-2777
US 1993-129204
           R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, MC, NL, PT, SE
                                                     PL 1992-301008 19920324 <--
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19931006 <--
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                                                                                    19950130 <--
                                                                                    19950602 <--
                                                                                    19950607 <--
                                                       SE 1991-1090 A 19910411 <--
SE 1993-2777 A 19930827 <--
US 1993-129204 B1 19931025 <--
US 1995-379471 B3 19950130 <--
US 1995-459660 A2 19950602 <--
US 1995-479494 A2 19950607 <--
PRIORITY APPLN. INFO.:
                                                       SE 1996-141 A 19960116 <--
CS 1993-2116 A 19920324 <--
EP 1992-907877 A3 19920324 <--
JP 1992-507195 A3 19920324 <--
WO 1992-SE186 A 19920324 <--
WO 1994-SE780 W 19940825 <--
AB
      There are described finely divided particles of a pharmaceutical
      substance, wherein the substance when submitted to water vapor gives off
      heat of less than 1.2 J per g, processes for their production and
      pharmaceutical formulations containing them. An example is given of
      salbutamol sulfate (25%) and lactose (75%) conditioned with water at
      relative humidity 55-65%, nonconditioned micronized substance mixture (5-8
      J/g) and conditioned micronized mixture (<0.5 J/g).
REFERENCE COUNT:
                                21
                                       THERE ARE 21 CITED REFERENCES AVAILABLE FOR THIS
                                       RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
      ANSWER 29 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
                            1999:42582 CAPLUS
DOCUMENT NUMBER:
                               130:100677
TITLE:
                              Antiasthmatic pharmaceutical composition containing
                              formoterol and rofleponide or their salts and
                               derivatives
INVENTOR(S):
                              Axelsson, Bengt; Kallstrom, Leif; Trofast, Jan
PATENT ASSIGNEE(S):
                              Astra Aktiebolag (Publ), Swed.
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PATENT NO. KIND DATE APPLICATION NO. DATE

WO 9900134 A1 19990107 WO 1998-SE1089 19980608 <-W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE,

PCT Int. Appl., 16 pp.

CODEN: PIXXD2

Patent English

SOURCE:

DOCUMENT TYPE:

PATENT INFORMATION:

FAMILY ACC. NUM. COUNT: 1

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DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP, KE, KG,
             KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX,
             NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT,
             UA, UG, UZ, VN, YU, ZW
         RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES,
             FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI,
             CM, GA, GN, ML, MR, NE, SN, TD, TG
     CA 2295076
                          Α1
                                19990107
                                            CA 1998-2295076
                                                                    19980608 <--
     AU 9881350
                          Α
                                19990119
                                            AU 1998-81350
                                                                    19980608 <--
     EP 1009408
                          Α1
                                20000621
                                            EP 1998-931163
                                                                    19980608 <--
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            AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
             IE, SI, LT, LV, FI, RO
     EE 9900594
                                 20000815
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                                                                    19980608 <--
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     TR 9903272
                          T2
                                 20000821
                                            TR 1999-3272
                                                                    19980608 <--
     BR 9810452
                          Α
                                20000905
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                                                                    19980608 <--
     HU 200002533
                          A2
                                20001228
                                            HU 2000-2533
                                                                    19980608 <--
     JP 2002510310
                          Т
                                20020402
                                            JP 1999-505479
                                                                    19980608 <--
     MX 9911676
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                                20000531
                                            MX 1999-11676
                                                                    19991214 <--
     NO 9906438
                          Α
                                20000228
                                            NO 1999-6438
                                                                    19991223 <--
PRIORITY APPLN. INFO.:
                                            US 1997-883823
                                                                 A 19970627 <--
                                            WO 1998-SE1089
                                                                 W 19980608 <--
     A composition or kit having as a first active ingredient formoterol
AB
     (I), or a salt or solvate derivative thereof, and having as a second active
     ingredient rofleponide (II), or a fatty acid ester thereof is disclosed.
     Also disclosed are methods for treating respiratory disorders using this
     composition or kit. II palmitate 10, dipalmitoylphosphatidylcholine 63,
     dimyristoylphosphatidylcholine 24, sodium dipalmitoylphosphatidylglycerol
     3, and racemic \alpha-tocopherol 0.1 parts were dissolved in 1300 parts
     tertiary butanol and the solution was freeze-dried to obtain a powder which
     was micronized to particle size of less than 5\mu m. I fumarate dihydrate
     0.5 parts was mixed with 79.5 parts of lactose monohydrate and micronized.
     This micronized mixture (80 parts) was added to the steroid/lipid
     freeze-dried powder (20 parts) and filled into a capsule for use in a dry
     powder inhaler.
REFERENCE COUNT:
                         5
                               THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS
                               RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
     ANSWER 30 OF 38
                      CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
                         1998:640189 CAPLUS
DOCUMENT NUMBER:
                         130:47320
TITLE:
                         Subsensitivity to bronchoprotection against adenosine
                         monophosphate challenge following regular once-daily
                         formoterol
                         Aziz, I.; Tan, K. S.; Hall, I. P.; Devlin, M. M.;
AUTHOR(S):
                         Lipworth, B. J.
CORPORATE SOURCE:
                         Dept of Clinical Pharmacology and Therapeutics,
                         Ninewells Hospital and, University of Dundee, Dundee,
SOURCE:
                         European Respiratory Journal (1998), 12(3),
                         580-584
                         CODEN: ERJOEI; ISSN: 0903-1936
PUBLISHER:
                         Munksgaard International Publishers Ltd.
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         English
     Regular treatment with inhaled long-acting \beta 2-agonists leads to
     subsensitivity to their bronchoprotective effects, although the effect of
     dosing frequency on this subsensitivity is not known. The aim of this
     study was to assess whether a once-daily dosing regimen with
     formoterol might be associated with a lesser degree of
     subsensitivity. In a randomized placebo-controlled double-blind,
     double-dummy crossover study 10 asthmatics treated with inhaled
     steroids (mean age 31 yrs, forced expiratory volume in one second
     (FEV1) 82% predicted) received 1 wk of treatment with: formoterol
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dry powder 24 µg twice daily (08:00 and 20:00 h); formoterol

24 μg once daily (20:00 h); or identical placebo. Adenosine monophosphate (AMP) bronchial challenge was performed 12 h after the first and the last dose of each treatment. There was significant loss of protection with formoterol twice daily between the first and last dose (geometric mean provocative concentration causing a 20% fall in FEV1 (PC20)): 475 vs. 129 mg $\bullet$ mL-1 (a 3.7-fold loss, p=0.006) and with formoterol once daily: 367 vs. 127 mg•mL-1 (a 2.9-fold loss, p=0.005), compared with placebo: 71 vs. 75 mg $\bullet$ mL-1 (nonsignificant). There was no significant difference in the degree of loss of protection between formoterol once and twice daily. For first-dose protection there was a significant difference between active treatments and placebo, but after the last dose the residual protection between active treatments and placebo was not significant. Thus, in patients taking inhaled corticosteroids, regular formoterol 24  $\mu g$  once daily induces a similar degree of subsensitivity to adenosine monophosphate bronchial challenge as with formoterol 24  $\mu g$ twice daily. This in turn suggests that even with a 24-h dosing interval there is the development of tolerance to formoterol by prolonged occupancy of airway  $\beta$ 2-adrenoceptors.

THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 17 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 31 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

1998:273803 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 129:63010

TITLE: Bronchodilators and corticosteroids in the treatment

of asthma

AUTHOR(S): Vianna, Elcio Oliveira; Martin, Richard J.

CORPORATE SOURCE: The Department of Medicine, National Jewish Medical

and Research Center, Denver, CO, 80206, USA

Drugs of Today (1998), 34(3), 203-223 CODEN: MDACAP; ISSN: 0025-7656 SOURCE:

PUBLISHER: J. R. Prous, S.A. DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

A review, with 164 refs. Despite advancements in treatment, the incidence of asthma, asthma-related deaths, and hospitalizations for asthma have increased significantly during the past decade. Although asthma mortality may now be decreasing, reasons for the worsening of morbidity and mortality in asthma remain unclear. These unexpected changes in asthma severity have sparked renewed interest in research into the pathogenesis and treatment of the condition.  $\beta$ 2-Adrenergic agonists are the most commonly used class of drugs for the treatment of asthma. Recent concerns about safety issues for  $\beta$ -agonists caused reevaluation of prescribing practices, and using them on an as-needed basis is now more frequently accepted and recommended. In acute asthma, a  $\beta 2$ -adrenergic agonist is still the medication of choice. Long-acting salmeterol and formoterol, administered only twice daily, can decrease symptoms of asthma during day and nighttime. On the other hand, the role of tolerance to their bronchodilator and bronchoprotective effects is still to be determined in the treatment of asthma. Theophylline, whose use has been limited by the potential for serious toxicity, may regain an important position in asthma treatment with the development of the knowledge about its anti-inflammatory actions. Dosing theophylline on a time-related basis also improves the risk/benefit ratio and makes it a useful drug for nocturnal asthma. Ipratropium bromide, an anticholinergic drug, still awaits a defined role in the treatment of asthma. Studies on its use for acute asthma have not achieved consensus and, for nocturnal asthma, the short duration of effect limits the benefits. Corticosteroids, including inhaled steroids, have measurable effects on symptoms, lung function, bronchial responsiveness, and inflammation associated with asthma. Side effects of chronic use limit systemic, but not inhaled administration. Newer prepns., like budesonide, flunisolide and fluticasone, decrease the incidence of possible side effects related to

inhaled steroids by having better ratio of topical to systemic potency. Daily doses <1600 µg of beclomethasone (or equivalent) are considered safe and higher doses should be reserved for patients with moderate to severe asthma. Although future trials are necessary to clarify many issues related to dosing of inhaled steroids, chronotherapy studies have shown that single administration between 3 and 5:30 p.m. may be as effective as 4 times a day dosing.

REFERENCE COUNT: 164 THERE ARE 164 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

**FORMAT** 

ANSWER 32 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1998:273646 CAPLUS

DOCUMENT NUMBER: 129:22771

TITLE: Long-acting inhaled  $\beta$ 2-agonists in asthma therapy Moore, Robert H.; Khan, Ayesha; Dickey, Burton F. AUTHOR(S): CORPORATE SOURCE: Baylor College of Medicine and the Houston Veterans

Affairs Medical Center, Houston, TX, 77030, USA

SOURCE: Chest (1998), 113(4), 1095-1108 CODEN: CHETBF; ISSN: 0012-3692

PUBLISHER: American College of Chest Physicians

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

A review with 183 refs. This paper reviews the pharmacol. of the long-acting inhaled  $\beta$ 2-agonists, salmeterol and formoterol, summarize results of their clin. trials, evaluate their safety records, and discuss their roles in the treatment of asthma. Preclin. and clin. studies involving salmeterol or formoterol were identified by a MEDLINE search, weekly computerized literature updates, and manual searches. Studies of satisfactory quality were chosen for review. Salmeterol and formoterol are potent and selective  $\beta$ 2-adrenoceptor agonists with durations of action >12 h. Their major differences are that formoterol has a rapid onset of action and is a partial agonist of high intrinsic efficacy, whereas salmeterol has a delayed onset and is a partial agonist of low intrinsic efficacy. Twice daily use of either drug results in improved lung function, reduced symptoms, and a better quality of life. These agents protect against exercise-induced asthma for 12 h and eliminate nighttime awakening in most patients. Limited tolerance develops, especially to their bronchoprotective effects, but their improvement of lung function is sustained. Regular use of salmeterol or formoterol provides subjective and objective amelioration of asthma in patients experiencing excessive symptoms or physiol. impairment despite the regular administration of low doses of inhaled corticosteroids (equivalent to approx. 500  $\mu$ g/d of beclomethasone). Intermittent use of either long-acting  $\beta 2$ -agonist can provide prolonged protection against exercise-induced asthma or nighttime symptoms. Patients should be instructed to continue taking inhaled steroids when long-acting  $\beta$ 2-agonists are administered on a regular schedule and to not take long-acting \$2-agonists between regularly scheduled doses. Used properly, they are effective and safe adjunctive agents in the treatment of asthma.

REFERENCE COUNT: 160 THERE ARE 160 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 33 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1998:160008 CAPLUS

DOCUMENT NUMBER: 128:238866

TITLE: Formoterol: an update of its pharmacological

properties and therapeutic efficacy in the management

of asthma

AUTHOR(S): Bartow, Rebecca A.; Brogden, Rex N.

Adis International Limited, Auckland, N. Z. CORPORATE SOURCE:

SOURCE: Drugs (1998), 55(2), 303-322 CODEN: DRUGAY; ISSN: 0012-6667

PUBLISHER: Adis International Ltd.
DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

A review with 89 refs. Formoterol, a selective β2-adrenoceptor agonist, produces ED-proportional bronchodilation, which persists for up to 12 h, in patients with reversible obstructive respiratory disease. Bronchodilation is significant within minutes of inhalation, maximal within 2 h, and at therapeutic doses is equivalent to that produced by standard doses of traditional  $\beta2$ -agonists. In single-dose studies comparing the two long-acting  $\beta$ 2-agonists formoterol and salmeterol, significant bronchodilation is achieved more rapidly with formoterol than salmeterol. Duration of bronchodilation is similar with both drugs. The therapeutic efficacy of inhaled formoterol has been equal to or greater than that of salbutamol (albuterol), fenoterol and terbutaline in both short and long term clin. trials. Formoterol reduces symptoms of nocturnal asthma and reduces the need for rescue medication compared with salbutamol. Recent studies have shown that the addition of inhaled formoterol 12 or 24µg twice daily to existing inhaled corticosteroid regimens improves lung function and reduces asthma symptoms compared with placebo. In one well designed study, the frequency of severe exacerbations of asthma over 12 mo was decreased by adding formoterol to existing regimens of inhaled corticosteroids. Tolerance to the bronchodilator response of formoterol has not been observed in long term clin. trials. Because of its long duration of action, formoterol offers significant therapeutic advantages over shorter-acting  $\beta$ 2-agonists in the treatment of nocturnal and exercise-induced asthma. Formoterol is effective in preventing exercise-induced asthma in adults and children and confers significantly more protection than salbutamol when administered 3 and 12 h before exercise. In general, inhaled formoterol is well tolerated. The most commonly reported adverse effects, tremor and palpitations, are those traditionally associated with the use of  $\beta$ 2-agonists. Oral formoterol and high doses of inhaled formoterol are associated with more adverse events than are the recommended doses of 6 to 24µq. Formoterol is currently recommended for use as an alternative to increasing inhaled steroid dosage in patients whose symptoms are inadequately controlled despite therapy with low to moderate doses of inhaled steroids and intermittent short-acting  $\beta$ 2-agonists, and results of recent studies support therapeutic guidelines. Long term clin. studies comparing formoterol and salmeterol have not yet been published. Further studies to evaluate the earlier use of formoterol in patients with mild to moderate asthma are needed to determine the role and long term safety of formoterol in the management of asthma.

REFERENCE COUNT: 89 THERE ARE 89 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L9 ANSWER 34 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1997:223895 CAPLUS

DOCUMENT NUMBER: 126:216649

TITLE: (Endo, syn) - (-) -3 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - 8 - (3 - Hydroxy - 1 - oxo - 2 - phenylpropoxy) - (3 - Hydroxy - 2 - phenylpropoxy) - (3 - Hydroxy - 2 - pheny

methyl-8-(methylethyl)-8-azoniabicyclo[3.2.1]octane

salts as antiasthmatic pharmaceuticals

INVENTOR(S): Banholzer, Rolf; Reichl, Richard; Disse, Bernd; Speck,

Georg

PATENT ASSIGNEE(S): Boehringer Ingelheim Kg, Germany

SOURCE: Ger. Offen., 5 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

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PATENT NO.
                        KIND DATE
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                                                                     DATE
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20010811 TW 1996-85108367
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                                  19970226 AU 1996-67397
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                       A 19981206 IL 1996-118986

A 19990202 BR 1996-9951

T 19990907 JP 1997-507251

C2 20010727 RU 1998-103895

T 20011115 AT 1996-927638

T 20020429 PT 1996-927638

B1 20020430 HR 1996-365

T3 20020516 ES 1996-927638
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B6
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                        B1 20021229 BG 1998-102202
A 19980130 NO 1998-424
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PRIORITY APPLN. INFO.:
                                              DE 1995-19528145
                                                                   A 19950801 <--
                                              WO 1996-EP3364
                                                                  W 19960731 <--
                                              US 1998-983420
                                                                   B1 19980114 <--
AΒ
     The use of title compds. as antiasthmatic pharmaceuticals is described.
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The use of title compds. as antiasthmatic pharmaceuticals is described. Thus, 18 g ipratropium bromide was purified by HPLC and resolved on Chiralcel OD columns to give both L and D-isomers. An aerosol formulation contained the L-isomer 0.005, sorbitan trioleate 0.1, and monofluorotrichloromethane and difluoromethane (2:3) to 100%.

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L9 ANSWER 35 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
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ACCESSION NUMBER:

1996:578310 CAPLUS

DOCUMENT NUMBER:

125:212342

TITLE:

A dose-response study with formoterol

Turbuhaler as maintenance therapy in asthmatic

patients

AUTHOR(S):

Schreurs, A. J. M.; Damste, H. E. J. Sinninghe; De

Graaff, C. S.; Greefhorst, A. P. M.

CORPORATE SOURCE:

Dept Pulmonology, Onze Lieve Vrouwe Gasthuis,

Amsterdam, 1090 HM, Neth.

SOURCE:

European Respiratory Journal (1996), 9(8),

1678-1683

CODEN: ERJOEI; ISSN: 0903-1936

PUBLISHER: Munksgaard
DOCUMENT TYPE: Journal
LANGUAGE: English

AB The aim of this randomized, double-blind, parallel group study was to determine

the lowest ED of 6, 12 and 24  $\mu g$  formoterol fumarate dihydrate Turbuhaler b.i.d. compared with placebo. The 4 wk treatment was preceded by a 1 wk run-in period. Morning peak expiratory flow (PEF) before intake of the study drug was the primary variable. Patients recorded PEF, prior to and 15 min after intake of the study drug (immediate response), asthma symptoms, and use of rescue medication morning and evening. Of 221 patients (71 females and 150 males), 194 were included in the efficacy per protocol (PP) anal.; mean age 47 yrs, mean forced expiratory volume in one second (FEV1) 2.01 L (58% of predicted), mean FEV1 reversibility 27% at entry. Ninety percent used inhaled steroids. Compared with placebo, 6  $\mu g$  formoterol b.i.d. was found to be the lowest ED in the morning (p=0.008) and evening (p=0.0041) PEF. The mean increases in PEF were 22 and 23 L\*min-1 resp., compared with placebo. After 6 µg formoterol, the mean immediate increase in morning PEF was 42 L\*min-1 compared to an increase of only 9 L\*min-1 after placebo (p<0.0001). All doses produced a statistically significant decrease in asthma symptoms, day and night, and the need for rescue medication at night. All doses were well-tolerated. In conclusion, the lowest ED in this study was formoterol Turbuhaler 6  $\mu$ g b.i.d.

L9 ANSWER 36 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1996:513756 CAPLUS

DOCUMENT NUMBER:

125:151185

TITLE:

Pharmaceutical aerosols containing sugars and

fluorocarbons or fluorochlorohydrocarbons

INVENTOR(S):

Green, Alexander Peter Glaxo Group Limited, UK

PATENT ASSIGNEE(S): SOURCE:

PCT Int. Appl., 25 pp.

CODEN: PIXXD2 Patent

DOCUMENT TYPE: LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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		3776						2006	0517										
	AT	1952	49			T			0815	_		995-					9951	222	<- <b>-</b>
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		5955						1999	0921	1	US 1	997-	8495	38		1	9970	624	<
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PRIC	RIT	APP	LN.	INFO	.:							994-				A 1	9941	224	<
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		1																	

AB Aerosol formulations for the administration of medicaments by inhalation comprises (a) particulate medicament; (b) at least one sugar; and (c) a fluorocarbon or hydrogen-containing chlorofluorocarbon propellant. Particulate lactose was dispensed into clean, dry glass bottles and the metering valve was fitted onto the bottles, then micronized fluticasone

propionate mixed with 1,1,1,2-tetrafluoroethane was pressure-filled into the canisters through the metering valve. The resultant inhalers delivered 25  $\mu g$  of fluticasone propionate/actuation.

L9 ANSWER 37 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1996:464566 CAPLUS

DOCUMENT NUMBER: 125:96168

TITLE: Propellant mixture for aerosol formulation INVENTOR(S): Sapsford, Andrew; Savage, Andrew Patrick

PATENT ASSIGNEE(S): Glaxo Group Limited, UK SOURCE: PCT Int. Appl., 19 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

P <i>i</i>	PATENT NO.					ND DATE			APPLICATION NO.						DATE			
W	9618	384			A1		1996	0620	,	WO 1	995-	EP48	24		1	9951	208 <	_
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ΙA	9642	604	•	•	Α		1996	0703		AU 1	996-	4260	4		1	9951	208 <	_
El	EP 789557				A1			0820	EP 1995-941077					19951208 <				
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J	2 1051			•	T			1013									208 <	
A	2165	75			Т		2002	0515		AT 1	995-	9410	77		1	9951	208 <	_
E.	3 2174	970			Т3		2002	1116		ES 1	995-	9410	77		1	9951	208 <	_
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_	6309	-						1030					83				829 <	
PRIORI'													0				210 <	
				• •									24				208 <	
	•									-			17		-		721 <	
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AB This invention relates to aerosol formulations which comprise (a) 1,1,1,2-tetrafluoroethane, 1,1,1,2,3,3,3-heptafluoropropane or mixts. thereof as propellant, (b) 1,1,2,2,3-pentafluoropropane as co-propellant, and (c) particulate medicament. A method of treating respiratory disorders which comprises administration by inhalation of an effective amount of a pharmaceutical aerosol formulation as defined is also described. Micronized salmeterol xinafoate was placed into a bottle with 1,1,2,2,3-pentafluoropropane and the bottle was sealed. 1,1,1,2-Tetrafluoroethane was added under pressure through the valve. The resultant inhaler delivered 25  $\mu g$  of salmeterol xinafoate per actuation.

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L9 ANSWER 38 OF 38 CAPLUS COPYRIGHT 2007 ACS on STN
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ACCESSION NUMBER: 1988:118997 CAPLUS

DOCUMENT NUMBER: 108:118997

TITLE: Compositions of liposomes and beta-2-receptor active

substances, for administration to the respiratory

tract

INVENTOR(S): Axelsson, Bengt Ingemar; Bystroem, Ulla Katarina;

Dahlbaeck, Carl Magnus Olof; Kaellstroem, Leif Arne;

Nilsson, Per Gunnar; Trofast, Jan William

PATENT ASSIGNEE(S): Draco AB, Swed.

SOURCE: PCT Int. Appl., 29 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PAT	PATENT NO.						KIND DATE				APPLICATION NO.					DATE		
WO	8705	803			A1	_	1987	1008		WO 1	987-	SE14	8		1:	9870:	323	<
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AU	8772	076			Α		1987	1020		AU 1	987-	7207	6		1:	9870:	323	<
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PRIORITY	APP	LN.	INFO	. :						SE 1	986-	1457			A 1:	9860	401	<
										WO 1	987-	SE14	8		A 1	9870	323	<

A pharmaceutical composition consists of a dry powder comprising liposomes and AΒ a  $\beta$ 2-receptor-active substance, the latter being preferably entrapped within the liposomes or portioned between the liposomes and an external phase. This composition is for administration to the respiratory trait, preferably by inhalation. Dipalmitoyl phosphatidylcholine 60 and cholesterol 60 mg dissolved in 10 g CHCl3 and 60 mg terbutaline sulfate dissolved in 1 mL H2O were emulsified, evaporated on a rotary evaporator to form a gel, and 3 g H2O added to the gel with mixing to form a liposome dispersion in which 38% of the terbutaline sulfate was encapsulated. Liposomes containing terbutaline sulfate were also tested for antiinflammatory and bronchospasmolytic effects (in rats and guinea pigs, resp.), with pos. results.

## **EAST Search History**

Ref # .	Hits	Search Query	DBs	Default Operator	Plurals	Time Stamp
L1	2074	formoterol	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	ON	2007/03/08 09:48
L2	2074	L1	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	ON	2007/03/08 09:49
L3	13366	"citrate buffer"	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	ON	2007/03/08 09:49
L4	43	L3 and L1	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	ON	2007/03/08 09:50
L5	4668601	aqueous or water	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	ON	2007/03/08 09:50
L6	1607	L1 and L5	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	ON	2007/03/08 09:50
L7 -	83788	steroid	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	ON	2007/03/08 09:50
L8	. 2977	fluticasone	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	ON	2007/03/08 09:51
L9	1104	L1 and L7	US-PGPUB; USPAT; USOCR; EPO; JPO; DERWENT	OR	ON	2007/03/08 09:51

## **EAST Search History**

L10	1273	L1 and L8	US-PGPUB; USPAT; USOCR;	OR	ON	2007/03/08 09:51
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